

1 stepped pressure equilibrium code : mp00ad

1. Solves Beltrami linear system for given helicity multiplier and poloidal flux, and returns an error function. Copy of mp00ac; planned redundant.
2. if `Lposdef=T`, the solution is provided by `F04ASF`, which assumes the matrix is symmetric positive-definite;
3. if `Lposdef=F`, the solution is provided by `F04ATF`;
4. The solution vector is “unpacked” by `up00aa`. The unpacking routine must be consistent with the “packing” description given in `global`.

1.0.1 error function

1. This routine returns an “error-function”, $\mathbf{F}(\mu, \delta\psi_p)$, defined as follows:
 - (a) if `Lconstraint.eq.0`, $\mathbf{F} = 0$.
 - (b) if `Lconstraint.eq.1`, $\mathbf{F}(\mu, \delta\psi_p) = (\mathbf{t}_{inn} - (p_{l-l} + \gamma p_{r-l})/(q_{l-l} + \gamma q_{r-l}), \mathbf{t}_{out} - (p_l + \gamma p_r)/(q_l + \gamma q_r))$, where, given the Beltrami field, the transform on the inner, \mathbf{t}_{inn} , and outer, \mathbf{t}_{out} , adjacent interfaces is computed by constructing straight-field line coordinates; the integers p_l , q_l , p_r and q_r are given on input; and $\gamma = (1 + \sqrt{5})/2$ is the golden mean.
 - (c) if `Lconstraint.eq.2`, $\mathbf{F}(\mu) = \int_l \mathbf{A} \cdot \mathbf{B} dv - \mathcal{K}_l$, where \mathcal{K}_l is the helicity given on input.

mp00ad.h last modified on 2012-05-01 ;
